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13. ABSTRACT (Maximum 200 Words)

In attempts to understand how the signaling by retinoic acid (the active vitamin A metabolite) is regulated we have been studying the retinoic acid binding protein called CRABP-II. These studies revealed that CRABP-II acts to enhance the transcriptional activities of RA and that it does so by directly delivering the hormone to its cognate transcription factor, RAR. Consequently CRABP-II dramatically sensitized cultured mammary carcinoma cells to RA-induced growth inhibition. Similarly, over-expression of CRABP-II inhibited mammary tumor growth in two different mouse models of cancer. CRABP-II may be a novel target for therapeutic and preventive strategies for retinoid-treatment of breast cancer. This project aims to delineate the mechanism by which CRABP-II modulates RA activity, especially as related to its ability to enhance the anti-proliferative action of the ligand. The first aim is to determine the extent to which CRABP-II acts in activating different isotypes of RAR. The second aim is to dissect the mechanism by which RA-induced growth inhibition is mediated. This will then allow for closer inspection of particular target genes that are under such control. The third aim is to understand the basis for RA-resistance of mammary carcinoma cells and how CRABP-II functions to overcome this resistance.

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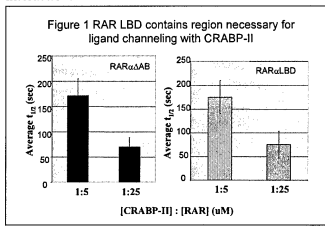
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Introduction

Retinoic acid (RA), the active metabolite of vitamin A, plays critical roles in embryonic development as well as growth and differentiation in adult mammals. Retinoic acid is currently used or is in clinical trials for treatment of a variety of different cancers, including breast cancer (1). Although retinoids are efficacious, pharmacological doses often result in toxicity (2). The ability of RA to induce growth arrest and apoptosis in carcinoma cells is mediated by the ligand-inducible transcription factors termed retinoic acid receptors (RARs). There are three isotypes of RAR, namely RARα, RARβ and RARγ (3). In cells, RA also associates with cellular retinoic acid binding proteins (CRABP-I and CRABP-II). We recently showed that CRABP-II carries RA from the cytoplasm to the nucleus where it channels the ligand directly to RARα via a transient protein-protein interaction (4, 5). This "ligand channeling" sensitizes cells to transcriptional activation by RAR (4). Indeed, over-expression of the binding protein dramatically lowered the effective RA concentration necessary to induce growth inhibition in mammary carcinoma cells (5). Similarly, over-expressing CRABP-II inhibited mammary tumor growth in two different mouse models of cancer. The goal of this project is to elucidate the mechanisms by which CRABP-II directs RA signaling to enhance anti-proliferative responses in mammary carcinoma cells.

Body Task 1. To determine which RAR isotypes is/are under CRABP-II regulation. Subtasks will overlap during this time period:

Our group has shown that CRABP-II channels RA directly to RARa via a transient protein-protein interaction (4). Theoretically, transfer of RA from CRABP to RAR may occur by one of two possible mechanisms: (1) RA may dissociate from the binding protein to the bulk aqueous phase prior to its association with RAR. (2) RA may transfer from CRABP to RAR by 'channeling', i.e. by a process that involves direct protein-protein interactions and that bypasses the bulk aqueous phase. The two pathways may be distinguished by the dependence of the rate constants of transfer of RA from the donor (CRABP) to the acceptor (RAR) on the concentration of the acceptor. If transfer follows mechanism (1), the rate-limiting step will be the dissociation of the ligand from CRABP, and the apparent rate constant will be independent of the concentration of RAR. On the other hand, if RA moves from CRABP to RAR by "channeling" (mechanism 2), then increasing the concentration of RAR (at a constant CRABP concentration), will result in a higher probability of productive donor-acceptor collisions, and thus, $t_{1/2}$ for the transfer reaction will become smaller as the acceptor concentration is raised. The observation that RA fluoresces much more when bound to CRABP-II versus when it is bound to RARa was used as a tool for measuring the transfer of ligand from binding protein to receptor (i.e. monitoring the loss of fluorescence of RA). The rate constants of transfer of RA from CRABP-I or CRABP-II to nearly full length RAR α (RAR $\alpha\Delta$ AB) were thus measured at different receptor concentrations. The rate of transfer from CRABP-II, but not CRABP-I, was found to strongly depend on the concentration of acceptor protein (4). Hence, CRABP-II (but not CRABP-I) delivers RA to RARα via direct protein-protein interaction.



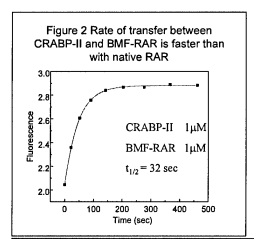
To determine the region of RAR α that mediates the interactions of the receptor with CRABP-II, we examined the ability of the ligand binding domain of RAR α (RAR α LBD) to engage in RA-channeling from the binding protein. Proteins were bacterially expressed and affinity purified. The dependence of the rate of ligand transfer from CRABP-II to the RAR LBD was then studied. As seen in figure 1, transfer was significantly facilitated upon increasing the acceptor/donor ratio from 1:5 to 1:25 with nearly identical kinetics of transfer as the nearly full length receptor. Hence, the LBD of the receptor contains the interaction domain for binding to CRABP-II and is

sufficient for ligand channeling.

In order to clarify which of the RAR isotypes is under direct control of CRABP-II, we planned to study the kinetic parameters of transfer of RA from CRABP-II to the three RAR isotypes. The ligand binding domains of RAR β and RAR γ were cloned into bacterial expression vectors and purified to near homogeneity. The fluorescence properties of retinoic acid bound to both RAR β and RAR γ were determined to ensure transfer experiments could be performed. Unfortunately, RA bound to these two isotypes fluoresces nearly as much as when bound to CRABP-II. Thus, transfer of RA from CRABP-II to RAR β and RAR γ could not be monitored by RA fluorescence.

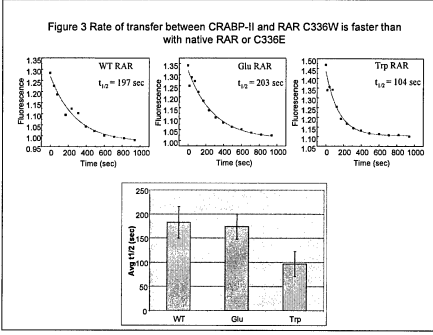
To overcome this setback, fluorescent probes can be used to label either the donor protein (CRABP-II) or acceptor protein (RAR). If ligand binding changes the fluorescence of the probe, this can be used as a measure of ligand transfer. A variety of fluorescent probes were tested for labeling CRABP-II. Some did not label the protein, some labeled the protein but ligand binding did not affect the fluorescence of the label, while others both labeled and responded to RA but did not channel with RARα. Thus, no suitable probe was found that could be used as a read-out for ligand transfer between CRABP-II and RAR.

Next, RARs were labeled with fluorescent probes. Again, a variety of probes were tested for labeling RARα. Labeling with the cycteine-attacking probe (5-bromo-methyl-fluorescein (BMF)) resulted in a labeled RARα whose fluorescence properties responded to ligand binding. Specifically, this protein displayed



fluorescence resonance energy transfer (FRET) between the RA and the protein-bound probe. Unfortunately, when RAR β and RAR γ were labeled with the same probe, the fluorescence of neither protein responded to ligand binding. Therefore, labeling RARs with BMF could not be used as a measure of ligand transfer between CRABP-II and RAR β and RAR γ .

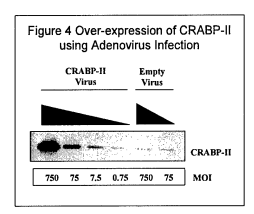
Channeling experiments were performed between CRABP-II and BMF-labeled RAR α . Surprisingly, RA channeled much faster to the labeled RAR α as compared to the unlabeled RAR α (Figure 2). RA transferred from CRABP-II to the labeled RAR with a $t_{1/2}$ of 36s, about 3-fold faster as compared with its rate of transfer to the native receptor. We hypothesize that RAR α was labeled near the site of interaction with CRABP-II and that the label facilitated ligand transfer.



With the help of mass spectrometry at the Harvard Microchemistry facility the labeling site was determined to be cysteine 336. Mutational analysis was performed to mimic the structure of the BMF at the site of labeling and tested for ligand transfer with CRABP-II. In these Cys336 was substituted with either Trp or Glu. As seen in figure 3 introducing a Trp in this location resulted in a protein that accepted RA from CRABP-II at a much faster rate than either the wild type or the Cys336Glu receptor. This indicates that (1) this region in RAR may be the domain responsible for the interaction with CRABP-II, and (2) that adding a bulky hydrophobic group at this position facilitates

ligand-channeling between these two proteins. We are now working in collaboration with Dr. Richard Gillilan to computationally "dock" the binding protein and receptor to determine if the region around Cys336 in RARa could be the interaction domain with CRABP-II.

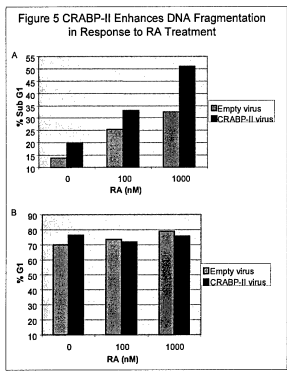
Task 2a. To determine the mechanism by which CRABP-II inhibits growth of mammary carcinoma cells.

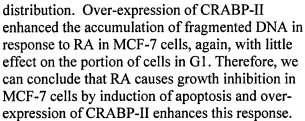


Our group has previously shown that over-expression of CRABP-II enhances the ability of RA to inhibit the growth of the mammary carcinoma cell line MCF-7 (5). The goal of this project is to determine the mechanism by which these cells are growth inhibited. We tested two signaling pathways that may respond to RA through RAR-mediated transcriptional regulation to induce growth arrest: (1) Apoptosis- RA may induce programmed cell death, characterized by cleavage of cytosolic cysteine proteases, caspases, and DNA fragmentation. (2) Cell cycle arrest-RA could induce growth arrest by blocking the cell cycle at a particular stage.

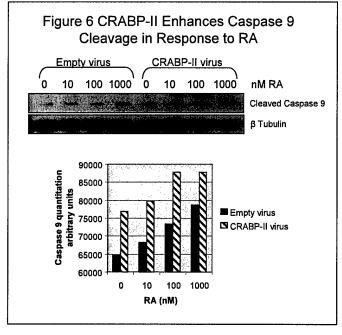
Apoptosis and cell cycle distribution were monitored following

RA treatment in MCF-7 cells that over-express CRABP-II compared to wild type cells. Over-expression of the binding protein in these cells was accomplished by viral infection with an adenovirus encoding the cDNA for CRABP-II (Figure 4). A multiplicity of infection (MOI) of 600 was used in subsequent experiments. Nuclei from MCF-7 cells treated with an empty virus (Ad0) or with the adenovirus harboring CRABP-II (Ad-CRABP-II) and treated with RA for 5 days were analyzed for DNA content by flow cytometry. In these experiments, a shift in percentage of G1 nuclei would indicate a cell cycle arrest while an increase in sub G1 population (fragmented DNA) would indicate the occurrence of programmed cell death. As seen in figure 5, retinoic acid treatment of MCF-7 cells caused an increase in the amount of fragmented DNA but had little effect on the cell cycle





Apoptotic responses are coordinated by a tightly regulated group of cysteine proteases named caspases. In a normal healthy cell, these proteins are found in an inactive state. Upon stimulation of apoptosis, procaspases are cleaved



to produce active proteases which then cleave cellular targets to propagate the apoptotic signal and induce death. Therefore, apoptotic responses can be detected by monitoring the appearance of the cleaved caspases. MCF-7 cells were treated with RA for 5 days and cell extracts were probed by western blotting for cleaved caspase 9 (Figure 6). Treatment of MCF-7 cells with RA caused a dose dependent activation of caspase 9. Cells that over-express CRABP-II show enhanced caspase 9 cleavage. Therefore, we conclude that RA-induced apoptosis in MCF-7 cells is a caspase 9 mediated event and over-expression of CRABP-II enhances the caspase response.

Task 2b. Effect of RA and CRABP-II on gene expression profiles in MCF-7 cells.

RA treatment causes growth inhibition in mammary carcinoma cells including MCF-7 cells. MCF-7 cells undergo apoptosis in response to RA treatment and this response is enhanced upon over-expression of CRABP-II. We would like to identify the genes involved in these responses. To this end, we carried out Affymetrix expression array analysis. In these, RNA is extracted from cells under particular assay conditions and corresponding cDNA is made with a fluorescent probe incorporated. This cDNA is then hybridized to a U133A chip containing single stranded DNA from most genes in the human genome, one gene at every spot on the chip. The labeled cDNA hybridizes to the spot containing its complimentary sequence. The more a given gene is expressed, the more signal will be measured from its spot on the chip. Thus, comparisons of gene expression can be made on an entire genome by comparing spot signal intensities from different conditions. We performed expression array studies using Affymetrix cDNA hybridization chips to identify genes that are upregulated in response to RA and also to over-expression of CRABP-II. Four conditions were analyzed by expression analysis, each one in triplicate; (1) empty virus, no RA treatment, (2) empty virus, 50 nM RA, 4 hours, (3) CRABP-II virus, no RA treatment, (4) CRABP-II virus, 50 nM RA, 4 hours. When MCF-7 cells were treated with RA, the expression of numerous genes was induced. Appendix 1 contains the entire list of genes induced by at least 1.32-fold in response to RA treatment. Contained in this list are many known RA target genes such as the RA degrading enzymes cytochrome P450s and homeobox genes. Importantly, the expression of many genes involved in apoptotic responses was also induced, as shown by the partial list of genes in Table 1. Specifically, caspase 7 and caspase 9 genes were both induced in response to RA. These observations support the experimental findings that RA triggers an apoptotic response in MCF-7 cells and suggest a mechanism by which the response is accomplished.

Table 1: Pro-Apoptotic Genes Induced by RA

Gene Name	Fold Change	P-Value
interferon regulatory factor 1	1.66	0.012061
tumor necrosis factor receptor superfamily, member 1A	1.65	3.99E-06
caspase 7, apoptosis-related cysteine protease	1.39	0.023309
caspase 9, apoptosis-related cysteine protease	1.37	0.012061
TIA1 cytotoxic granule-associated RNA binding protein	1.33	0.027382
small inducible cytokine subfamily E, member 1	1.32	0.067476
interleukin 6 signal transducer (gp130, oncostatin M receptor)	1.32	0.067476

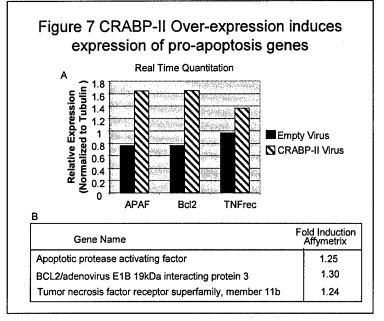
The known function of CRABP-II is to bind RA and deliver it to RAR in the nucleus thereby enhancing the transcriptional activity of the receptor. This binding protein has no known function in the absence of RA. Surprisingly, the expression array screens indicated that a number of pro-apoptotic genes are induced by simply over-expressing CRABP-II in the absence of RA treatment (Figure 7B). Among these are 3 genes that we chose to use in order to validate the Affymetrix data, namely: Apoptosis protease activating factor, BCL2/adenovirus E1b 19kDa interacting protein 3, and tumor necrosis factor receptor superfamily, member 11b. For validation studies, quantitative real-time PCR (Q-PCR) was carried out (Figure 7A). cDNA from the array was subjected to Q-PCR using Taqman probes. Primer and probe sets for all genes tested were obtained from Applied Biosystems as Assays on Demand. The relative standard curve method was used to quantitate the relative expression for each gene. Fold induction was calculated by dividing the expression of each gene in the CRABP-

II over-expressing cDNA by the expression of that gene in the empty virus treated cDNA. As seen in figure 7, all three genes were found to be induced in samples from CRABP-II over-expressing cells. The Q-PCR thus verified the validity of the observations derived from the Affymetrix array screen. This up-regulation of pro-

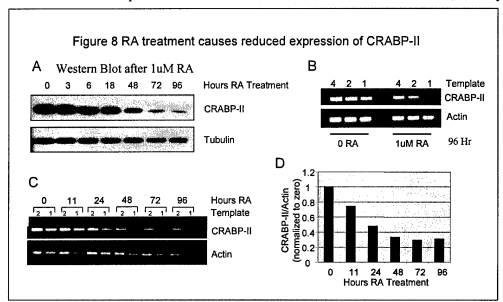
apoptotic genes may explain how, in the previous assays, over-expression of CRABP-II in the absence of RA caused an increase in all the apoptotic markers tested. Importantly, this may point to a novel, RA-independent function of CRABP-II. Our current working hypothesis is that CRABP-II may heighten cellular responses not only to RA signaling via RAR but also to a variety of other apoptotic signals. We are currently testing the effect of over-expression of CRABP-II on the effectiveness of other known apoptosis inducers such as etoposide or TNFα.

Task 3. Understand the underlying basis for RA resistance in mammary carcinoma cells

A frequent complication in utilizing RA for cancer therapy is the development of



RA-resistance in tumors. Understanding the mechanisms that underlie RA-resistance in cancer cells are thus of significant clinical importance. We have observed that mammary carcinoma cells that over-express CRABP-II become more sensitive to RA-induce growth inhibition. Conversely, we find that cells that do not express CRABP-II are resistant to this anti-proliferative effect. Therefore, we examined the effect of RA on the expression of CRABP-II. The data reveal that RA treatment resulted in a decrease in CRABP-II protein in MCF-7 cells (Figure 8A). This effect may originate from two possible responses: (1) RA treatment may lead to a degradation of CRABP-II protein, or (2) RA treatment may result in a decrease in mRNA levels. To determine whether this loss of protein is due to a decrease in CRABP-II mRNA level, semi-quantitative PCR was



performed on extracts from cells treated with RA. Figure 8B-D shows that the CRABP-II RNA level in MCF-7 cells is down regulated in response to RA treatment. RA-induced down-regulation of CRABP-II expression may comprise an important feature through which carcinoma cells are rendered RA-resistant. On going studies aim to understand the molecular mechanisms underlying down regulation of CRABP-II expression.

Key Research Accomplishments

- The ligand binding domain of RARa is sufficient for interaction and ligand channeling with CRABP-II
- The region around Cys336 is RARα is the potential interaction region with CRABP-II necessary for channeling
- Retinoic acid causes apoptosis in MCF-7 cells with little effect on cell cycle distribution.
- Over-expression of CRABP-II in MCF-7 cells enhances the apoptotic response to RA.
- Expression array data indicate that expression of certain pro-apoptotic genes is induced upon treatment with RA
- Over-expression of CRABP-II in the absence of RA causes an increase in pro-apoptotic genes indicating a novel ligand-independent function for this binding protein.
- RA-resistance of carcinoma cells may stem from RA-induced down-regulation of CRABP-II.

Reportable Outcomes:

There are no reportable outcomes to this date.

Conclusions:

RA is currently used or is in clinical trials for therapy of a variety of cancers, however, at pharmacological doses it is often toxic. Our lab is investigating approaches that will allow for sensitization of cancer cells to RA chemotherapy in order to increase the therapeutic efficacy of this compound. We have found that a RA binding protein (CRABP-II) functions to inhibit mammary carcinoma cell proliferation in culture as well as tumor progression *in vivo*. Therefore, CRABP-II may be a novel target for therapeutic and preventive strategies for treatment of breast cancer. The mechanisms by which this binding protein acts to modulate RA signaling and enhance the anti-proliferative activities of this compound in breast cancer are being studied. CRABP-II enhances RA-induced apoptosis in MCF-7 cell. Interestingly, CRABP-II appears to function by two separate mechanisms. The protein enhances RA-induced apoptosis and it also increases expression of pro-apoptotic genes even in the absence of RA. These observations suggest that CRABP-II possesses a novel ligand-independent function which contributes to its anti-carcinogenic activity. Studies carried out during this funding period also revealed that RA down-regulates the expression of CRABP-II in carcinoma cells. This activity may be an important factor in the development of RA-resistance in carcinomas.

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- 5. A. S. Budhu, N. Noy, *Mol Cell Biol* **22**, 2632-41 (Apr, 2002).

 UniGene Name 9 cytochrome P450, family 26, subfamily A, polypeptide 1 4 cytochrome P450 retinoid metabolizing protein 9 homeo box A5 1 short-chain dehydrogenase/reductase 1 8 homeo box A1 				1 pleckstrin homology domain containing, family E (with leucine rich repeats) member 1 8 regulator of G-protein signalling 16 7 cytochrome P450, family 24, subfamily A, polypeptide 1 6 retinoic acid induced 3 5 Homo sapiens, clone IMAGE:4816940, mRNA 4 endothelial PAS domain protein 1 5 sordium channel proportion and a labba	
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kallikrein 6 (neurosin, zyme) c-Mpl binding protein hypothetical protein, estradiol-induced	interferon regulatory factor 1 snermidine/snermine N1-acetyltransferase	tumor necrosis factor receptor superfamily, member 1A	GATA binding protein 6	v-maf musculoaponeurotic fibrosarcoma oncogene homolog F (avian) forkhead hov A1	ankyrin repeat domain 3	FLN29 gene product	serine threonine kinase 39 (STE20/SPS1 homolog, yeast)	degenerative spermatocyte homolog, lipid desaturase (Drosophila)	intercellular adhesion molecule 1 (CD54), human rhinovirus receptor	selectin L (lymphocyte adhesion molecule 1)	TATA box binding protein (TBP)-associated factor, RNA polymerase I, A, 48kDa	WW domain-containing protein 1	protein tyrosine phosphatase, receptor type, E	chromosome 6 open reading frame 62	solute carrier family 31 (copper transporters), member 2	kinesin family member 23	pro-oncosis receptor inducing membrane injury gene	pre-B-cell leukemia transcription factor 1	Nedd4 binding protein 3	transmembrane protease, serine 4	TBC1 domain family, member 4	chromosome 20 open reading frame 45	F-box only protein 34	neural precursor cell expressed, developmentally down-regulated 9	frizzled homolog 7 (Drosophila)	lipocalin 2 (oncogene 24p3)	basic helix-loop-helix domain containing, class B, 2	protein kinase C, delta	dual-specificity tyrosine-(Y)-phosphorylation regulated kinase 2	prostaglandin E synthase	lysophospholipase I
0.75 0.75 0.75	0.73	0.72	0.72	0.72	0.7	0.7	0.69	0.69	0.68	0.68	0.68	0.68	0.68	0.68	0.67	0.67	0.67	0.67	0.67	0.67	99.0	99'0	0.66	0.65	0.65	0.65	0.64	0.64	0.64	0.64	0.64
204733_at 214155_s_at 218245_at	202531_at 203455_s_at	207643_s_at	210002_at	36711_at 204667_at	221215_s_at	35254_at	202786_at	209250_at	202638_s_at	204563_at	206613_s_at	212637_s_at	221840_at	222309_at	204204_at	204709_s_at	211967_at	212148_at	214775_at	218960_at	203386_at	217851_s_at	218539_at	202149_at	203706_s_at	212531_at	201170_s_at	202545_at	202968_s_at	210367_s_at	212449_s_at

t calpastatin	4 chromosome 10 open reading frame 2	3 core-binding factor, runt domain, alpha subunit 2; translocated to, 3	3 LIM and cysteine-rich domains 1	2 KIAA1609 protein	1 glutamate-ammonia ligase (glutamine synthase)	5 cystatin A (stefin A)	5 solute carrier family 22 (organic cation transporter), member 4	3 hypothetical protein MGC10940	3 Rab coupling protein	3 prostate differentiation factor	9 C-terminal binding protein 2	9 LSM5 homolog, U6 small nuclear RNA associated (S. cerevisiae)	9 PTPL1-associated RhoGAP 1	3 mannosyl (alpha-1,6-)-glycoprotein beta-1,2-N-acetylglucosaminyltransferase	8 TBC1 domain family, member 8 (with GRAM domain)	8 retinoic acid induced 17	3 forkhead box C1	8 KIAA1609 protein	7 neuroepithelial cell transforming gene 1	7 spinocerebellar ataxia 1 (olivopontocerebellar ataxia 1, autosomal dominant, ataxin 1)	7 karyopherin (importin) beta 3	7 kidney ankyrin repeat-containing protein	7 tumor endothelial marker 6	7 activated RNA polymerase II transcription cofactor 4	5 candidate mediator of the p53-dependent G2 arrest	5 hypothetical protein MGC2776	5 cullin 4A	5 restin (Reed-Steinberg cell-expressed intermediate filament-associated protein)	5 hairy and enhancer of split 1, (Drosophila)	5 ELOVL family member 6, elongation of long chain fatty acids (FEN1/Elo2, SUR4/Elo3-like, yeast)	5 transmembrane protein 5	5 hypothetical protein from clone 643	5 inhibitor of Bruton's tyrsoine kinase	5 nucleoporin 160kDa	5 spastic ataxia of Charlevoix-Saguenay (sacsin)
0.64	0.64	0.63	0.63	0.62	0.61	9.0	9.0	9.0	9.0	9.0	0.59	0.59	0.59	0.58	0.58	0.58	0.58	0.58	0.57	0.57	0.57	0.57	0.57	0.57	0.56	0.56	0.55	0.55	0.55	0.55	0.55	0.55	0.55	0.55	0.55
212586 at	218590_at	208056_s_at	218574_s_at	221843_s_at	215001_s_at	204971_at	205896_at	209512_at	219681_s_at	221577_x_at	201218_at	202904_s_at	203910_at	203102_s_at	204526_s_at	212124_at	213260_at	65438_at	201829_at	203232_s_at	211953_s_at	213005_s_at	217853_at	221727_at	219370_at	219581_at	201424_s_at	201975_at	203394_s_at	204256_at	204807_at	209679_s_at	210970_s_at	212709_at	213262_at

# N = #	0.54 epithelial V-like antigen 1 0.54 likely ortholog of mouse Sh3 domain YSC-like 1 0.54 transcription factor 8 (represses interleukin 2 expression)	0.54 · down-regulator of transcription 1, 1 br-binding (negative cofactor z) 0.54 · vascular endothelial growth factor 0.54 · v-crk sarcoma virus CT10 oncogene homolog (avian)-like 0.54 · raft-linking protein	0.54 zinc finger protein 3 (A8-51) 0.54 KIAA0648 protein 0.54 pleckstrin homology-like domain, family A, member 1 0.54 family with sequence similarity 18 member B	- V N	0 10 2 0 - 15	 0.53 hypothetical protein STRALL11499 0.53 mitochondrial folate transporter/carrier 0.52 A kinase (PRKA) anchor protein 1 0.52 TGFB1-induced anti-apoptotic factor 1 0.52 serine (or cysteine) proteinase inhibitor, clade A (alpha-1 antiproteinase, antitrypsin), member 3 0.52 interferon gamma receptor 1 	
218605_at 219540_at 201830_s_at 203705_s_at	203780_at 204019_s_at 208078_s_at	210510_at 210512_s_at 212180_at 212646_at	212684_at 213984_at 217996_at 218446_s_at	218696_at 219178_at 221193_s_at 201219_at	203100_s_at 205083_at 212522_at 213139_at 214771_x_at 216268_s_at	218251_at 221020_s_at 201675_at 202039_at 202376_at 202776_s at	202837_at 207935_s_at 212060_at 212249_at 213304_at

213704_at 218538_s_at 218889_at 222361_at 39891_at 202345_s_at 203217_s_at 203422_at 204700_x_at 209422_at 201467_x_at 201467_x_at 201701_s_at 201701_	0.52 0.52 0.52 0.52 0.52 0.51 0.51 0.51 0.52 0.53 0.53 0.54 0.54 0.54 0.54 0.54 0.54 0.54 0.54	Rab geranylgeranyltransferase, beta subunit MRS2-like, magnesium homeostasis factor (S. cerevisiae) AD24 protein similar to beta-tubulin 4Q similar to beta-tubulin 4Q similar to beta-tubulin 4Q similar to beta-tubulin 4Q similar to beta-tubulin 3G clary acid binding protein 5 (psoriasis-associated) hypothetical protein MCC28875 chromosome 20 open reading frame 104 UDP-N-acetyl-alpha-D-galactosamine:polypeptide N-acetylgalactosaminyltransferase 7 (GalNAc-T7) stimulated by retinoic acid gene 6 calpastatin splicing factor, arginine/serine-rich 3 NK3 transcription factor related, locus 1 (Drosophila) Rho interacting protein 3 dehydrogenase/reductase (SDR family) member 2 hypothetical protein MGC28875 epitheilal protein fost in neoplasm beta protein kinase C, eta serologically defined colon cancer antigen 16 tight junction protein 3 (zona occludens 3) nucleolar GTPase laminin, alpha 3 homeo box 85 homeo box 86 homeo box
218947_s_at 220890_s_at 203563_at 203759_at 207181_s_at	0.49 0.48 0.48 0.48 0.48	hypothetical protein FLJ10486 DEAD (Asp-Glu-Ala-Asp) box polypeptide 47 actin filament associated protein sialyltransferase 4C (beta-galactoside alpha-2,3-sialyltransferase) caspase 7, apoptosis-related cysteine protease
208754_s_at 214658_at	0.48 0.48	nucleosome assembly protein 1-like 1 CGI-109 protein

 0.48 tripartite motif-containing 31 0.48 Homo sapiens cDNA FLJ13695 fis, clone PLACE2000124. 0.48 DKFZP586N0721 protein 0.48 hypothetical protein FLJ10199 0.48 egl nine homolog 1 (C. elegans) 0.48 sec13-like protein 	1.47 protein tyrosine phosphatase, non-receptor type 12 1.47 transcription elongation regulator 1 (CA150) 1.47 KIAA0372 gene product 1.47 component of oligomeric golgi complex 2 1.48 scription associated phosphoprotein 2 1.49 ubiquitin D 1.40 nuclear receptor co-repressor 2 1.40 nuclear receptor co-repressor 2 1.40 microtubule-associated protein 1 light chain 3 beta 1.40 cDC-like kinase 1 1.41 hypothetical protein FLJ10493 1.47 WD repeat domain 3		 0.46 zinc finger protein 23 (KOX 16) 0.46 keratin 23 (histone deacetylase inducible) 0.46 chromosome 14 open reading frame 58 0.46 Fas apoptotic inhibitory molecule 0.45 RIO kinase 3 (yeast) 0.45 transcriptional co-activator with PDZ-binding motif (TAZ) 0.45 phosphatidylserine decarboxylase 0.45 G protein-coupled receptor, family C, group 5, member B
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215444_s_at 216159_s_at 218284_at 218815_s_at 221497_x_at 221931_s_at	202006_at 202396_at 2033049_s_at 203073_at 204362_at 205890_s_at 207760_s_at 208766_s_at 208766_s_at 208776_s_at 218683_s_at 218683_s_at 218683_s_at 218772_x_at	220161_s_at 221648_s_at 202307_s_at 204341_at 206935_at 208802_at 209498_at 211764_s_at 213280_at	213934_s_at 218963_s_at 219316_s_at 220643_s_at 202130_at 202133_at 202392_s_at 203632_s_at

203984_s_at 204175_at 204244_s_at 204244_s_at 204244_s_at 20495_s_at 205618_at 205618_at 208407_s_at 209451_at 218319_at 218319_at 20193_at 200776_s_at 200776_s_at 200776_s_at 20193_at 200776_s_at 20193_at 20193_at 202704_at 202704_at 202704_at 202704_at 202704_at 202706_at 202704_at 202706_at 204521_at 205479_s_at 204521_at 204521_at 204521_at 204521_at 204521_at 204521_at 204521_at 205479_s_at 205479_s_at 212499_s_at 212499_s_at 214240_at 214240_at 214240_at 214240_at 214240_at 214240_at 21433_at 219443_at 219443_at	0.0.0.0.0.0.0.0.0.0.0.0.0.0.0.0.0.0.0.	caspase 9, apoptosis-related cysteine protease zinc finger protein activator of S phase kinase DKFZP434H132 protein pre-B-cell leukemia transcription factor 1 proline-rich Gla (G-carboxyglutamic acid) polypeptide 1 catenin (cadherin-associated protein), delta 1 TRAF family member-associated NFKB activator N-acylsphingosine amidohydrolase (acid ceramidase) 1 pellino homolog 1 (Drosophila) hypothetical protein FLJ20360 lactamase, beta 2 basic leucine zipper and W2 domains 1 fatty-acid-Coenzyme A ligase, long-chain 2 transducer of ERBB2, 1 hairy and enhancer of split 1, (Drosophila) protein predicted by clone 23733 MAD, mothers against decapentaplegic homolog 3 (Drosophila) protein predicted by clone 23733 MAD, mothers against decapentaplegic homolog 3 (Drosophila) plasminogen activator, urokinase Ste20-related serine/threonine kinase serine (or cysteine) proteinase inhibitor, clade B (ovalbumin), member 1 myosin IB chromosome 14 open reading frame 32 homeo box A10 galanin solute carrier family 38, member 6 G protein-coupled receptor 48 pleckstrin homology domain containing, family F (with FYVE domain) member 2 homeo box C13
	0.43 0.43 0.43 0.43 0.43	nomeo box C13 aryl hydrocarbon receptor hypothetical protein HSPC111 phosphoribosyl pyrophosphate synthetase 2 LIM domain only 2 (rhombotin-like 1) phospholipase A2, group X

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201038_s_at 201448_at 202350_s_at	0.41 0.41 0.41	hypothetical gene supported by U60823; U73477; X75090; AF025684; AY007110; BC007200; NM_006305 TIA1 cytotoxic granule-associated RNA binding protein matrilin 2
	0.41	sparc/osteonectin, cwcv and kazal-like domains proteoglycan (testican)
	0.41	acyl-Coenzyme A oxidase 3, pristanoyl
204656_at	0.41	SHB (Src homology 2 domain containing) adaptor protein B
	0.41	hematopoietically expressed homeobox
	0.41	hypothetical protein MGC29875
	0.41	nudix (nucleoside diphosphate linked moiety X)-type motif 4
	0.41	EH-domain containing 1
	0.41	sterol carrier protein 2
	0.41	H3 histone, family 3B (H3.3B)
	0.41	mitochondrial tumor suppressor gene 1
	0.41	nudix (nucleoside diphosphate linked moiety X)-type motif 4
	0.41	KIAA0582 protein
	0.41	regulator of chromosome condensation (RCC1) and BTB (POZ) domain containing protein 1
	0.41	hypothetical protein FLJ20508
	0.41	lysophosphatidic acid phosphatase
	0.41	opsin 3 (encephalopsin, panopsin)
	0.41	nuclear receptor coactivator 6 interacting protein
	0.41	nudix (nucleoside diphosphate linked moiety X)-type motif 6
	0.41	RAD51-like 3 (S. cerevisiae)
	0.41	Homo sapiens mRNA; cDNA DKFZp762M127 (from clone DKFZp762M127)
200608_s_at	0.4	RAD21 homolog (S. pombe)
201129_at	0.4	splicing factor, arginine/serine-rich 7, 35kDa
201410_at	0.4	pleckstrin homology domain containing, family B (evectins) member 2
201599_at	0.4	ornithine aminotransferase (gyrate atrophy)
202541_at	0.4	small inducible cytokine subfamily E, member 1 (endothelial monocyte-activating)
203098_at	0.4	chromodomain protein, Y chromosome-like
203124_s_at	0.4	solute carrier family 11 (proton-coupled divalent metal ion transporters), member 2
203693 <u>s</u> at	0.4	E2F transcription factor 3
203745_at	0.4	holocytochrome c synthase (cytochrome c heme-lyase)
204137_at	0.4	transmembrane 7 superfamily member 1 (upregulated in kidney)
204342_at	0.4	calcium-binding transporter
204354_at	0.4	protection of telomeres 1

205158_at 205434_s_at 205681_at 205681_at 206770_s_at 208322_s_at 209101_at 212195_at 21325_at 21325_at 213626_at 213792_s_at 213792_s_at 213792_s_at 218188_s_at 218158_s_at 218158_s_at	0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0	 ribonuclease, RNase A family, 4 AP2 associated kinase 1 0.4 AP2 associated kinase 1 0.6 BCL2-related protein A1 0.6 Solute carrier family 35 (UDP-N-acetylglucosamine (UDP-ClcNAc) transporter), member A3 0.7 solute carrier family 35 (UDP-N-acetylglucosamine (UDP-ClcNAc) transporter), member A3 0.8 sialyltransferase 4A (beta-galactoside alpha-2,3-sialyltransferase) 0.9 connective tissue growth factor 0.1 interleukin 6 signal transducer (gp130, oncostatin M receptor) 0.2 phosphatidylinositol binding clathrin assembly protein 0.3 phosphatidylinositol binding clathrin associated, actin dependent regulator of chromatin, subfamily a, member 5 0.4 hypothetical protein 1 0.5 NS1-associated protein 1 insulin receptor 1 insulin receptor 2 NS1-associated protein 3 adaptor protein containing pH domain, PTB domain and leucine zipper motif 3 adaptor protein FLJ20485 4 hypothetical protein FLJ20485
221478_at	0.4	BCL2/adenovirus E1B 19kDa interacting protein 3-like
43511_s_at	4.0	Homo sapiens mRNA; cDNA DKFZp762M127 (from clone DKFZp762M127)